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# A multi-task learning approach for prediction of missing bioactivity values of compounds for the SLC transporter superfamily

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Solute carrier (SLC) transporters constitute the largest family of membrane transport proteins in humans. They facilitate the movement of ions, neurotransmitters, nutrients, and drugs. Given their critical role in regulating cellular physiology, they are important therapeutic targets for neurological and psychological disorders, metabolic diseases, and cancer. Inhibition of SLC transporters can modulate substrate gradients, restrict the cellular uptake of nutrients and drugs, and thereby facilitate specific pharmacological effects. Despite their pharmaceutical relevance, many SLC transporters remain understudied. Having a complete bioactivity matrix of associated compounds can expand the knowledge base of SLC ligands, enlarge the information pool to guide downstream processes, and promote informed decision-making steps in the discovery of new drug candidates for SLC transporters. To address the data sparsity of available compound-bioactivity values causing inhibitory responses for SLC transporters, we employed a multi-task learning (MTL) approach with a data imputation objective. By leveraging relationships between related tasks, deep learning has previously shown promise in imputing compound bioactivities across multiple assays. We developed a multi-task deep neural network (MT-DNN) to predict and impute missing pChEMBL ( $-\log(\text{IC}_{50})$ ) values across the SLC transporter superfamily. With a data matrix density of 2.53% and an  $R^2$  of 0.74, our model demonstrated robust predictive performance. Specifically, we predicted missing values for 9122 unique compounds across 54 SLC targets spanning various folds and subfamilies, generating 480 133 predictions from 12 455 known interactions. The advantages of the multi-task learning approach were indicated in the ability of certain targets to leverage the shared representation of knowledge and acquire increased predictive accuracy over single-task learning (STL) counterparts. Despite the limitations set by low data density, activity cliffs, and inter-protein heterogeneity, the MT-DNN showed promising potential as a tool to address data sparsity within the SLC superfamily.

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## Introduction

SLC transporters are one of the largest groups of proteins in humans, accounting for 30% of the total proteome. The SLC superfamily consists of more than 450 members that are organized into 65 families defined by sequence homology and physiological functions.<sup>1</sup> As membrane-bound proteins, SLC members transport different types of molecules across the membranes. These include amino acids, lipids, sugars, ions, neurotransmitters, and drugs. They exhibit different underlying transport dynamics by utilizing ion concentration gradients as symporters, translocating substrates in the opposite direction of ions as antiporters, and channel-like properties for single transported molecules as uniporters. SLC transporters are distributed across different tissues and organs, including the brain, liver, and kidneys. Being involved in the uptake and efflux

of molecules relevant to phase I and phase II metabolism, they impact the absorption and elimination of drugs. Gene mutations in some SLC proteins that lead to altered or impaired ability to transport endogenous compounds contribute to the development of neurological diseases, cholesterol/bile transport defects, and cancer.<sup>2,3</sup> Inhibition of SLC transporters can facilitate pharmacological effects with therapeutic purposes intended for treatment of different diseases.<sup>4</sup> Targeting sugar transporters of the SLC5 family to modulate substrate translocation has been proposed for treatment of diabetes, cancer, and cardiovascular diseases.<sup>5</sup> Transport of neurotransmitters by SLC6 family members can be disrupted by inhibitors that bind to the transporter and block the conformational changes required for substrate uptake. Such inhibition regulates neurological responses and forms the basis of several therapeutic strategies, including the treatment of depression, attention-deficit hyperactivity disorder (ADHD), and other neuropsychiatric conditions.<sup>6</sup> Organic anion transporters (OATs) of the SLC22 family mediate the transport of negatively

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charged molecules such as nutrients, metabolites, toxins, and drugs. Inhibitors of OATs can modulate substrate uptake and excretion, thereby influencing the absorption and renal clearance of co-administered drugs and ultimately prolonging their exposure in patients.<sup>7</sup> Conversely, organic cation transporters (OCTs) of the same family mediate the uptake of cationic molecules in different tissues. Inhibition of OCT3 increases extracellular levels of serotonin and norepinephrine, thereby providing an alternative therapeutic approach for the treatment of depressive disorders. However, the scarcity of specific OCT3 ligands continues to limit therapeutic development.<sup>8</sup> Concomitant inhibition of OCTs and SLC47 members, such as multi-drug and toxin extrusion protein 1 (MATE1), has been recognized as clinically relevant due to their involvement in drug–drug interactions. Successful efforts have been made with *in silico* methods to predict the inhibitory activity of MATE1 and aid experts in the drug discovery process.<sup>9</sup>

Increased utility of artificial intelligence in drug discovery has followed technological advancements over the past years.<sup>10</sup> Qualitative and quantitative properties of compiled datasets have a major impact on the performance of machine learning and deep learning models. The data used for training can contain various proportions of missing information due to technical errors or the intrinsic nature of the objective.<sup>11</sup> This is evident in healthcare cases where patient information is incorrectly documented or entirely missing.<sup>12</sup> In the context of pharmaceutical research, bioactivity information of compounds can be scarce depending on the target.<sup>13</sup> This becomes evident when comparing industry data warehouses with public data repositories such as ChEMBL.<sup>14,15</sup> Alternatively, there is less interest in targets that cause rare diseases, which leads to a limited information pool available on their ligands.<sup>16</sup> Moreover, investigative studies showed that bioactivity errors and dataset size directly influence accuracy scores of machine learning models.<sup>17</sup>

Data imputation is a statistical method that is used to replace the missing values. In the context of pharmacoinformatics, it utilizes sparsely filled experimental data to impute the bioactivities or properties of compounds by leveraging the relationships between available datapoints.<sup>18</sup> Several data imputation methods have been proposed in the cases of biological assays with sparse data matrices: (i) a single-task DNN model that was trained on a set of compounds to predict pIC50 values for each assay independently,<sup>19</sup> (ii) a multi-task deep neural network that was trained to predict the bioactivity values across each assay simultaneously,<sup>20</sup> (iii) feature net approach with two training steps that involve predicting activity values for each task independently in the first step and using the predicted values as features together with compound descriptors to retrain the model in the second step,<sup>19</sup> (iv) the Alechmite DNN that uses compound descriptors and activity values as inputs; the missing activity values are substituted for the mean values, and the model is trained to iteratively update the predictions until no further improvement is observed.<sup>18,21</sup> Additional relevant approaches to consider for data imputation are pQSAR and matrix factorization methods like Macau.<sup>22,23</sup> The common feature among most imputation

approaches that outperform single-task QSAR methods is that they establish the relationships between the tasks as endpoints to facilitate some form of knowledge sharing or transfer.<sup>24</sup> This can lead to increased performance in sparse datasets, extend the domain of applicability for dissimilar molecules, and save computation time.<sup>25</sup>

MTL aims to generate the shared representation of knowledge by simultaneous training on multiple data domains from different tasks and, thereby, improve generalization capabilities.<sup>26</sup> This approach can be employed to predict the bioactivity values or properties of compounds.<sup>27</sup> Rather than focusing on activity predictions for a single protein, MTL can facilitate predictions with multiple outputs and benefit from establishing relationships between related tasks. Besides having the advantage of natural regularization, MTL models can transfer information between different tasks and increase overall accuracy. This is specifically relevant in the cases of individual proteins with scarce data, as they can potentially benefit from other similar proteins through shared knowledge. To avoid confusing the model, it is important to consider degrees of similarity and how correlated the tasks (proteins) are.<sup>28</sup>

In this study we explore MTL as a prediction tool with a data imputation objective to replace the missing pChEMBL values of 9122 unique compounds across 54 protein members of the SLC superfamily as individual tasks. We developed two MT-DNNs that were trained on different types of descriptors for comparative purposes. In addition to the overall performance, target-based performance was evaluated to inspect the abilities of individual proteins to exploit the advantages of an MTL approach. We highlight challenges associated with activity cliffs and employ dimensionality reduction methods to analyze discrepancies in the chemical space between high- and low-scoring targets. Finally, we explore comparative analysis between prediction and imputation abilities of the model. Even though protein targets belong to a single superfamily, degrees of heterology between them can vary depending on the organizational hierarchy they occupy. They belong to different families, subgroups, and folds, and have different transport mechanisms, substrates, and tissue localizations.<sup>29</sup> Because of this, it is important to consider putative inconsistencies in degrees of correlation between the targets as individual tasks that can lead to negative transfers.<sup>30</sup>

## Materials and methods

### Data acquisition and processing

All data operations and experiments were conducted using the Python programming language (v3.11.5). The list of SLC transporters was acquired from the Resolute Knowledgebase<sup>31</sup> and merged with the UniProt human-protein dataset<sup>32</sup> to associate UniProt-ID values with the corresponding target names. The compiled SLC protein dataset contained 446 unique UniProt-ID numbers, gene names, and protein names. The list of UniProt-IDs was queried *via* the ChEMBL Database API (ChEMBL web services) to retrieve the corresponding ChEMBL-IDs of individual targets.<sup>33</sup>



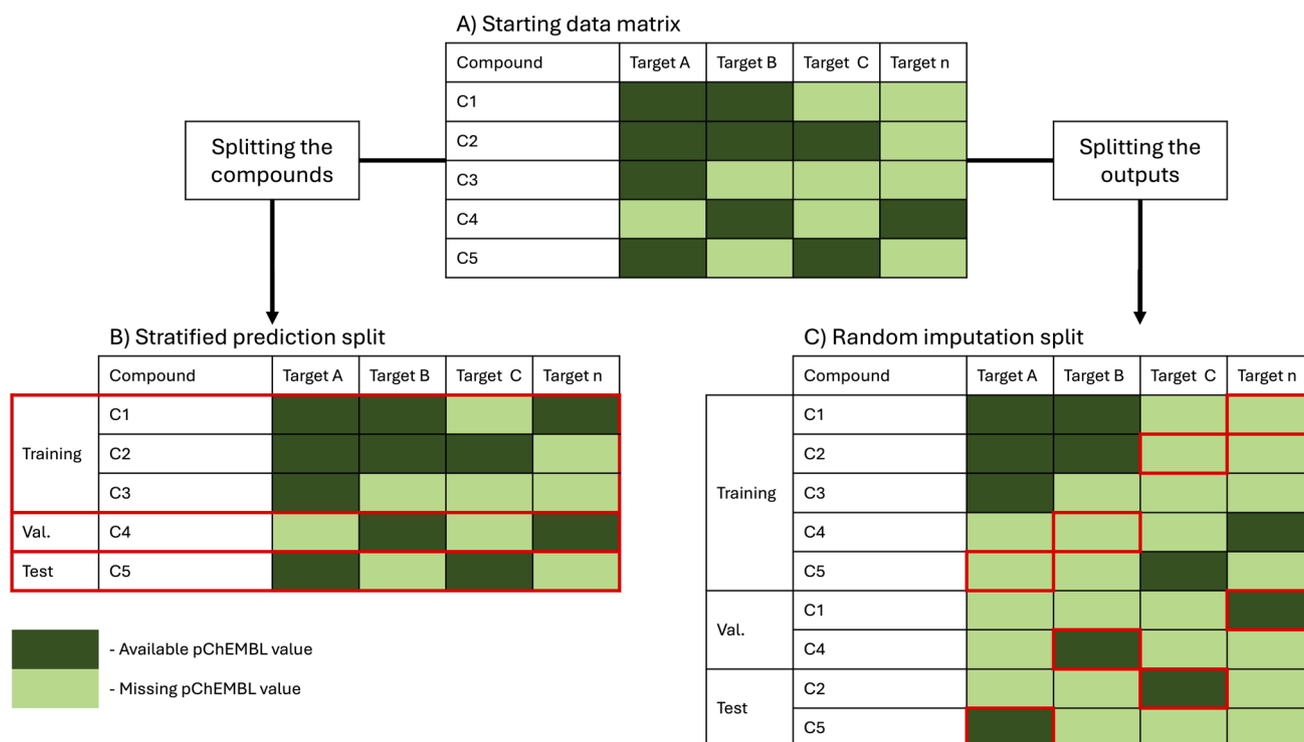
Table 1 Quantitative properties of the dataset after processing

Families	Targets	Compounds	Data density
25	54	9122	2.528%

The ChEMBL-IDs of targets were imported into a Jupyter Notebook for compound retrieval from the ChEMBL33 database and processing with RDKit (v2024.03.05).<sup>14</sup> Compounds that contained inorganic elements were excluded, stereochemistry information was removed, and standardized SMILES strings were calculated. The lowest pChEMBL value in the dataset was 4.0, corresponding to an IC<sub>50</sub> of 100 000 nM. During an analysis, it could not be confirmed whether these entries at the lower end of the inhibitory potency distribution represented actual dose–response measurements or simply reflected minimum reporting thresholds (100 000 or 10 000 nM). Therefore, data points with IC<sub>50</sub> values of exactly 100 000 and 10 000 nM were excluded. To ensure consistency across the 54 modeled targets, entries containing the substrings “muta” or “recombi” in the assay description column were removed. Mutagenized or recombinant variants could introduce inconsistencies in target representation and confound the learning process of the multi-task model. Excluding these entries helped maintain a uniform definition of targets and improved the

reliability of cross-target bioactivity estimation. Duplicate measurements of pChEMBL values from identical compound–target pairs were replaced by the mean values if the SEM (standard error of the mean) was lower than 0.2. Conversely, compound–target pairs with an SEM higher than 0.2 were excluded from the dataset. Remaining duplicate entries were consequently removed so that only one measurement was kept per compound–target pair in the dataset.

After the processing steps, the table was formatted into a data matrix intended for a multi-task imputation problem. Each compound was assigned the name of the associated target in the label column designated for the stratified split. In cases where multiple targets were measured for a single compound, the name of the target with the lowest number of compounds in the dataset was assigned. This ensured proportional representation of each target in the training, validation, and test sets. A split ratio of 80 : 10 : 10 for training, validation, and test sets was maintained across all targets to ensure consistency. To establish valid evaluation across all endpoints, targets with fewer than ten compounds were excluded so that each modeled target contained at least one datapoint in both validation and test sets. Finally, the data matrix comprised 9122 unique compounds across 54 targets and 25 families, with a data density of 2.53% (Table 1). Two sets of descriptors were used for comparative purposes. The Continuous Data Driven Descriptors (CDDD) are



**Fig. 1** Simplified representation of the two data-splitting approaches. (A) Mock bioactivity data matrix showing five unique compounds across four different targets. Dark green cells indicate available bioactivity values, whereas light green cells represent missing values. (B) Stratified prediction split: entire data points (*i.e.*, compounds with their complete bioactivity profiles) are allocated to the training, validation, or test sets according to targets as labels. (C) Random imputation split: compounds with more than two available bioactivity measurements are split into two data points with differing bioactivity profiles. One data point is assigned to the training set and the other is randomly allocated to either the validation or test set.



Table 2 Data splitting

Dataset	Stratified prediction split	Random imputation split
Training	7297	9122
Validation	913	1200
Test	912	1200

calculated using an autoencoder model that was trained to represent molecular formats such as SMILES as vectors of 512 continuous values.<sup>34</sup> The repository containing the model and instructions to calculate CDDs is available on GitHub (<https://github.com/jrwnter/cddd>). A second set of descriptors was 1024 extended connectivity fingerprints with a radius of 3 (ECFP6), which were calculated with a CDK module.<sup>35</sup> The ECFP6 are circular fingerprints that contain structural information of a given molecule in the form of the binary values (0/1).<sup>36</sup>

Two different approaches, shown in Fig. 1, were chosen to split the data. (Fig. 1B) Stratified prediction split: a target-based stratified split was used to allocate 80% of the compounds to the training set and 10% to the validation and test sets, respectively. Each compound, along with its complete bioactivity profile, was treated as a single data point to ensure non-overlapping compound distributions across sets. Random imputation split (Fig. 1C): in this approach, data were split according to bioactivity measurements rather than individual compounds. Compounds with two or more available bioactivity values were divided into two data points, each representing a different subset of their bioactivity profiles. One data point was assigned to the training set, and the other was randomly allocated to either the validation or test set. This procedure artificially introduced missing values in the training data, which were then used for evaluation. Consequently, all compounds from the validation and test sets were also present in the training set but with differing bioactivity profiles. This setup was designed for imputation-specific evaluation and contrasts with the compound-based prediction split due to intentional data leakage. Although identical compounds were used during training, different output values were used to evaluate knowledge sharing/transfer across targets.

Quantitative profiles after applying two splitting approaches are shown in Table 2.

### Hyperparameter optimization

Hyperparameters were tuned with the Optuna library (v3.6.1) by setting the range of values or categories for the model type, quantity of hidden layers, hidden-layer sizes, sizes of task-specific layers, learning rate, and dropout rate.<sup>37</sup> Parameters in Table 3 are shown as ranges of values in “()” and categories as “[]”. The Bayesian search determined the optimal parameters by minimizing the MSE value of the validation set calculated at the end of each training iteration (trial). A task-specific domain with two layers was conditioned with the model type being set to “shared and specific”. If the model type was set to “shared”, the task-specific layer took the role of an output layer with the size corresponding to the number of modeled tasks (targets).

Table 3 Hyperparameter ranges and categories set for the Bayesian search of the MT-DNN models

Parameter	Suggested value range-[]/category-()
Model type	(Shared, shared and specific)
Hidden depth	(2, 3)
1st hidden layer	[100–1200]
2nd hidden layer	[50–500]
3rd hidden layer	[20–200]
1st specific layer	[3–30]
2nd specific layer	[2–50]
Learning rate	[0.000040–0.000170]
Dropout rate	[0.07–0.50]
Patience	[7–10]

Analogously, the size of the third layer was conditioned by the Hidden depth being set to 3. The batch size was set to 128. After 300 iterations, the best-scoring set of parameters was selected to determine the architecture and hyperparameters of the model.

### MT-DNN architecture

The models were built as multiple-input/multiple-output feed-forward backpropagation deep neural networks with PyTorch (v2.3.0). They consist of an input layer, two or three fully connected hidden layers, and one task-specific/output layer. LeakyReLU was set as an activation function. The sizes of the input and output layers were determined by the number of descriptors and tasks, respectively. The sizes of hidden layers were determined through hyperparameter optimization steps. The loss function was calculated as the masked mean squared error (MSE) between the predicted and true values in the output layer (1). To ensure that the model parameters were exclusively updated with the MSE/loss derived from available values during training, predicted and true outputs were masked at indices where no bioactivity measurement was originally observed. Adam was selected as a method for stochastic optimization of the models. The MT-DNNs were trained on the training set with the validation MSE/loss calculated at each epoch to evaluate generalization performance and learning dynamics. To limit overfitting, a failsafe mechanism, referred to previously as the patience parameter, was set to terminate the training loop at the step where no improvement of the mean MSE validation value was observed for 7–10 consecutive epochs.

$$\text{MSE (masked)} = \frac{\sum_{i=1}^n m_i (y_i - \hat{y}_i)^2}{\sum_{i=1}^n m_i}$$

### Model evaluation

The model was evaluated with descriptor-type performance discrepancies, overall-prediction performance, learning dynamics, outlier vs. inlier evaluation, target-based prediction performance, visualization of the chemical space, and overall-imputation performance with benchmarking. To assess how well the entire dataset was represented by the model, 5-fold CV was implemented for the prediction splits. Every metric indicated in Table 4 followed the masked indexing principle of the



**Table 4** Evaluation metrics for performance assessment of the regression models

Metric	Formula
Mean squared error	$\text{MSE} = \frac{1}{n} \sum_{i=1}^n (y_i - \hat{y}_i)^2 \quad (2)$
Coefficient of determination	$R^2 = \frac{\sum_{i=1}^n (y_i - \hat{y}_i)^2}{\sum_{i=1}^n (y_i - \bar{y})^2} \quad (3)$
Cross-validated (CV) $R^2$	$Q^2 = R^2(\text{CV}) \quad (4)$
Model training score	$\text{MTS} = R^2(\text{training}) \quad (5)$
Mean absolute error	$\text{MAE} = \frac{1}{n} \sum_{i=1}^n  y_i - \hat{y}_i  \quad (6)$
Root mean squared error	$\text{RMSE} = \sqrt{\frac{\sum_{i=1}^n (y_i - \hat{y}_i)^2}{n}} \quad (7)$

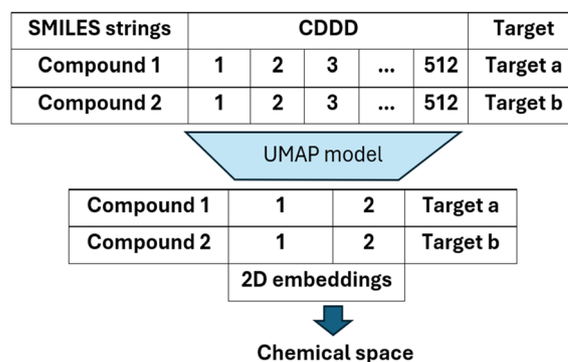
MSE/loss function (1). Modules for data splitting, CV, and evaluation were adopted from scikit-learn.<sup>38</sup> Three categories were chosen for an MTL-STL comparative analysis with aggregate task-specific evaluation: (1) unweighted: calculating mean scores and percentages by assuming that all targets are equally relevant. (2) Weighted: mean scores and percentages are adjusted by assigning higher relevance to targets with a higher number of compounds in the dataset. (3) Inversely weighted: mean scores and percentages are adjusted by assigning higher relevance to targets with a lower number of compounds in the dataset. In this evaluation step, category 3 could be considered the most representative form of quality assessment for the knowledge sharing capabilities of the multi-task imputation framework, indicating advantages/disadvantages the model has for underrepresented targets. Three targets that represent distinct performance profiles observed during target-based evaluation were further analyzed with 2D chemical space representation. Furthermore, as outlined in the section Data acquisition and processing, conceptionally different approaches for data splitting were applied: (i) in the case of the stratified prediction split, compounds are split such that entire molecules are held out from training and the task corresponds to out-of-sample prediction. In this setting, the model is evaluated on unseen compounds, and the objective is to assess generalization performance, descriptor suitability, and target-specific predictive accuracy. (ii) In random imputation split, the same MT-DNN architecture was retrained under a matrix-completion setting, where the compound set is fixed and bioactivity values for certain targets are intentionally removed during training. In this case, compounds are present as input entities during training but removed target labels do not contribute to the MSE/loss function. Model performance is then evaluated on these withheld entries. This setting corresponds to a data imputation scenario, *i.e.*, estimating missing values within a partially observed compound–target activity matrix.

## Single-task models

Single-task deep neural networks (ST-DNN) and single-task random forest (ST-RF) models were generated with PyTorch and scikit-learn respectively to conduct a comparative analysis between the multi-task and single-task learning approaches. The hyperparameters for each single-task model were tuned with a Bayesian search. The hyperparameter ranges shown in SI Table 1 for the ST-DNN, as well as the model architectures, were set to reflect the MT-DNN model. The hyperparameter ranges of the ST-RF models are shown in SI Table 2. To maintain consistency across all three approaches, the optimum hyperparameters were selected based on the minimum MSE value over 300 trials with Optuna. The data were split randomly, and 5-fold CV was implemented in the evaluation step to assess the performance across the entire dataset.

## Chemical space

The multi-dimensional information of chemical descriptors is reduced to two or three dimensions for visual interpretation. This way, relationships between different compounds can be observed while taking into consideration the context of the entire chemical dataset. Dimensionality reduction algorithms can utilize different types of chemical descriptors to project the chemical space from a set of compounds.<sup>39</sup> The principles behind the Uniform Manifold Approximation and Projection (UMAP) are a series of mathematical operations that reduce the dimensionality of features/descriptors while preserving the local structure of data. This method attempts to keep neighboring data points in close proximity and thereby preserve their relationship when mapping from high dimensionality in low-dimensional space. UMAP (v0.5.5) was used for visual representation of the chemical space in this study.<sup>40</sup> A set of 9122 unique compounds with 512 CDDD descriptors was used for the chemical space analysis. After the dimensionality reduction (Fig. 2), 2D embeddings were associated with the corresponding SMILES strings. The scatterplot of the chemical space was generated with the Matplotlib library. Compounds were color-coded based on the associated targets that were selected in the evaluation. The parameters shown in SI Table 3 for the



**Fig. 2** Chart showing the dimensionality reduction of 512 CDDD descriptors into 2D embeddings for an exemplary set of 2 compounds intended for chemical space visualization.



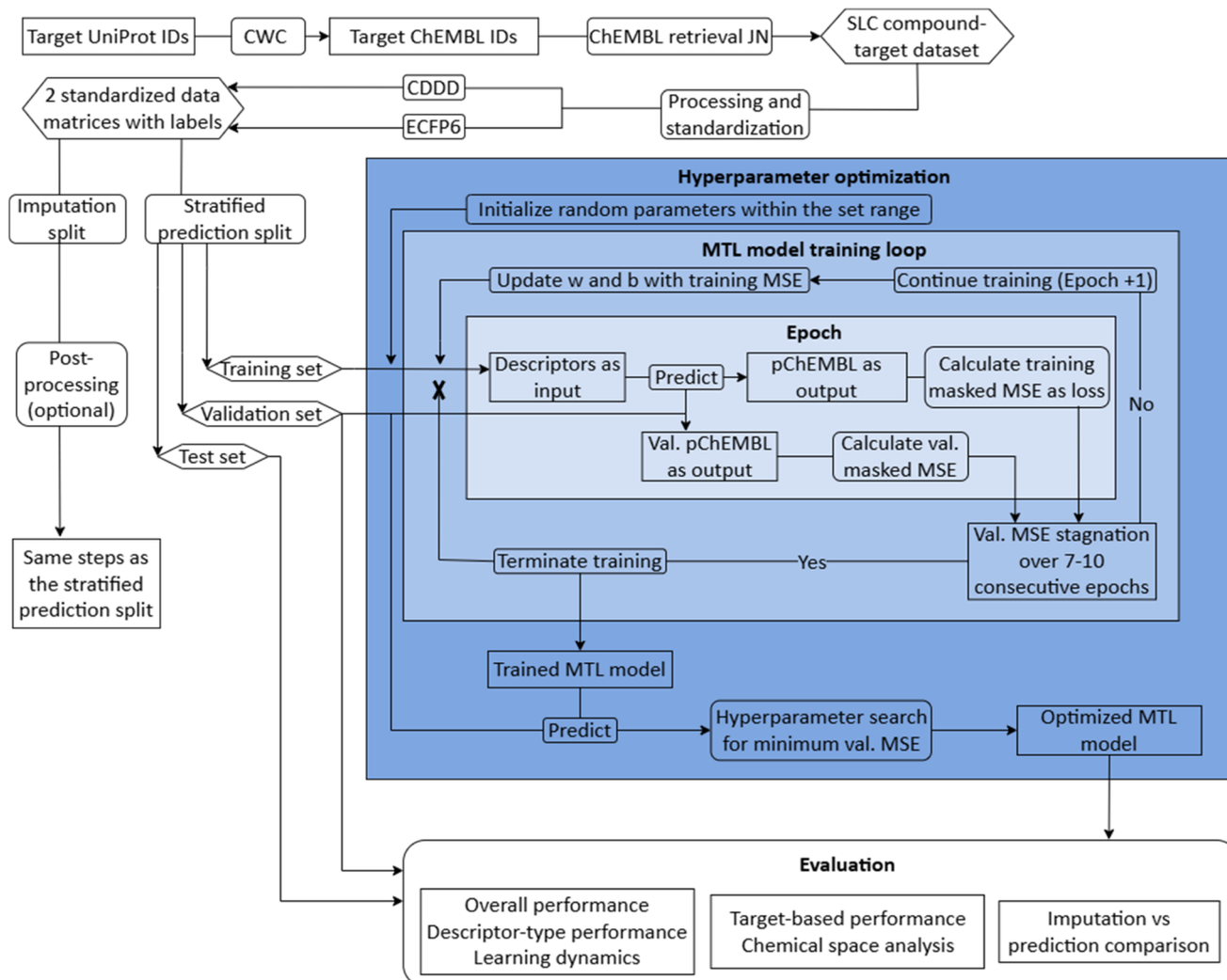


Fig. 3 Overview of the most relevant steps in the development of the MT-DNN for bioactivity prediction and data imputation evaluation. Starting from the top left point, the workflow depicts steps that were necessary in retrieving the data, processing data, calculating descriptors, performing data splits, developing, and evaluating the model. Blue squares highlight the steps specifically related to the training and hyperparameter optimization of the model, with a breakdown of a single epoch in the center. The workflow chart was created at <https://app.diagrams.net/>.

UMAP model were tuned to represent the compounds belonging to common families as neighbors in 2D and thereby form family-associated clusters.

### Data imputation models for benchmarking

To evaluate the imputation performance of the MT-DNN, six additional imputation frameworks were adopted in this workflow for comparative analysis. The choice of benchmarking models was based on accessibility and technical reasons related to time and cost restraints. Three models were selected from the scikit-learn library (SimpleImputer, KNNImputer, and IterativeImputer) and three from the fancyimpute library (SoftImpute, IterativeSVD, and MatrixFactorization).<sup>41</sup> Datasets obtained through the imputation split (see Fig. 1) were used to develop the models. To ensure consistency in the comparative analysis, the same data format, comprising CDDD descriptors and a sparsely populated bioactivity matrix used for MT-DNN training, was also employed to train the benchmarking

imputation models. Models followed a similar strategy to the MT-DNN development by using training data for fitting, validation data to optimize hyperparameters, and test data for evaluation. Parameter ranges shown in SI Table 4 were set for a grid search of the respective imputation models.

For easier comprehension of the project development pipeline, an overview of the entire workflow can be seen in Fig. 3.

## Results and discussion

### MT-DNN prediction performance

The parameters and architecture of the models shown in Table 5 were selected according to a Bayesian search performed with the Optuna library. Both models achieved the lowest MSE value of the validation set without the task-specific domain, indicating that the standalone shared domain of the hidden layers contributes positively to the overall performance of the model. Furthermore, the CDDD model has 2 hidden layers, whereas the



Table 5 Optimized hyperparameters for CDDD and ECFP6 models<sup>a</sup>

Hyperparameters	Models	
	CDDD	ECFP6
Input size	512	1024
HL1 size	800	1160
HL2 size	250	230
HL3 size	—	190
Task-specific size	54	54
Learning rate	0.000043	0.000046
Dropout rate	0.1	0.08
Patience	10	7

<sup>a</sup> Hidden layer (HL).

ECFP6 model has 3 hidden layers in the shared domain. Considering the results from the evaluation metrics in Table 6, the CDDD model was able to achieve higher performance with less architectural complexity than the ECFP6 model. With  $Q^2$  and  $R^2$  values being closer to the MTS in Table 6, the CDDD model indicates a lower tendency to overfit to the training data. Additionally, the learning dynamics comparison in Fig. 4 shows that the MSE loss of the validation set from the CDDD model has a closer loss curve to the training data when compared to the ECFP6 model. This trend remains consistent overall and was a decisive factor for continuing to use CDDD descriptors in further development and evaluation of the MTL approach.

Descriptor-associated discrepancy in the performance proposes higher ability by CDDD representation of compounds to capture the diversity of chemical space that was necessary to model the SLC superfamily. As only one additional descriptor type was considered, further comparative studies would be required for conclusive statements.

### Activity cliffs

Analysis of pChEMBL predictions with the highest error values (HEV) for a given compound–target pair was conducted to identify the shortcomings of the MTL approach with CDDD descriptors. The SLC6A4 target (serotonin transporter, SERT) had instances of both HEV and low error value (LEV) predictions that showcase putative effects of activity cliffs on the performance of the model. Both examples are compared with their nearest neighbors identified *via* the NearestNeighbors module from scikit-learn and corresponding activity profiles. The HEV case in Table 7 shows that the predicted pChEMBL of 6.39 falls within the range of values of the nearest neighbors for the same target. However, the actual pChEMBL value of the HEV compound is 4.0. Considering that the

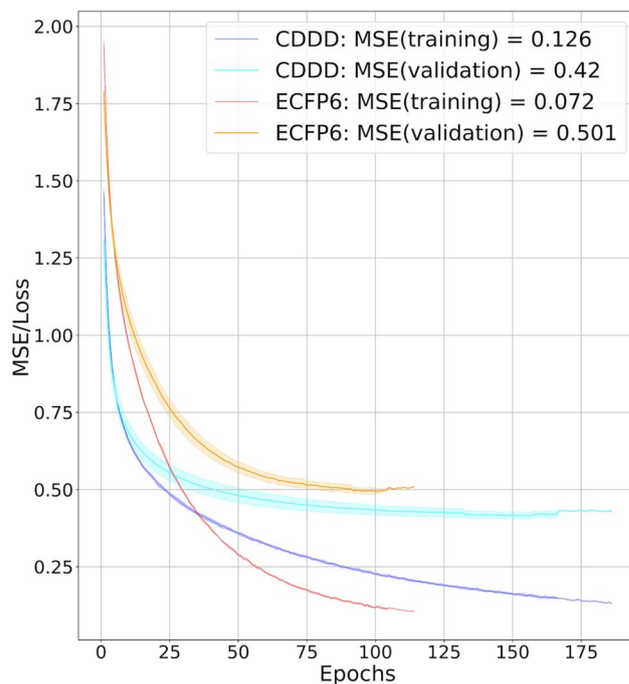


Fig. 4 Learning dynamics graph showing the mean values of MSE/loss over epochs for the training and validation data. The values were derived from the 5-fold CV to represent the learning dynamics across the entire dataset. MSE/loss values are shown on the y-axis across epochs represented on the x-axis.

three closest neighbors of the HEV compound in the training set had a mean pChEMBL value of 6.8, overprediction by the model could be attributed to the training-set bias. Inversely, the LEV case in Table 7 has the actual and predicted values of the test set compound falling within the bioactivity ranges of the nearest neighbors in the training set. When considering further comparison of the two examples, a higher degree of structural similarity between the LEV compound and its nearest neighbors can be observed. The challenge associated with activity cliffs could be addressed by implementing proportional distribution of biological activities across similar compounds or respective targets. This approach could ensure sufficient representation of various activity ranges between different targets and induce a balanced learning profile by the model.

### Target-based evaluation

Performance scores for individual targets were calculated as  $Q^2$  values using MT-DNN, ST-DNN, and ST-RF models during 5-fold

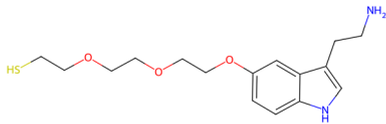
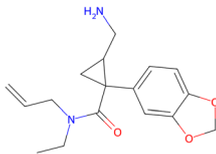
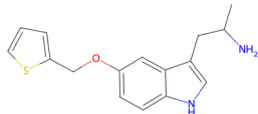
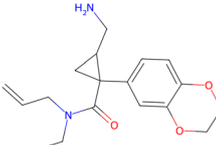
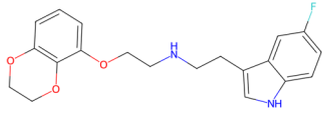
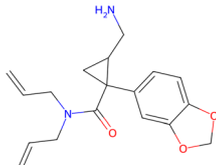
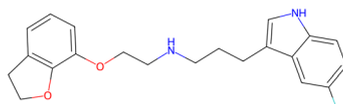
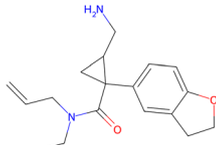
Table 6 Evaluation results for CDDD and ECFP6 models<sup>a</sup>

Model	MTS↑	Cross-validation				Test data			
		$Q^2$ ↑	MSE↓	MAE↓	RMSE↓	$R^2$ ↑	MSE↓	MAE↓	RMSE↓
CDDD	0.915	<b>0.716</b>	<b>0.420</b>	<b>0.486</b>	<b>0.650</b>	<b>0.739</b>	<b>0.362</b>	<b>0.451</b>	<b>0.602</b>
ECFP6	<b>0.952</b>	0.661	0.501	0.536	0.708	0.689	0.432	0.477	0.657

<sup>a</sup> Bold – the best value in a row.



Table 7 HEV and LEV comparison for the SLC6A4 target<sup>a</sup>

HEV compound (activity cliff example) – left panel	Actual pChEMBL	Predicted pChEMBL	LEV compound – right panel	Actual pChEMBL	Predicted pChEMBL
CHEMBL264262 	4.0	6.39	CHEMBL406789 	6.82	6.82
<b>Nearest neighbors of the HEV compound in the training set</b>			<b>Nearest neighbors the LEV compound in the training set</b>		
CHEMBL1255834 	5.32	—	CHEMBL427904 	7.19	—
CHEMBL124700 	7.07	—	CHEMBL259694 	7.24	—
CHEMBL338982 	8.01	—	CHEMBL258180 	6.37	—

<sup>a</sup> ChEMBL-IDs and actual pChEMBL values are annotated for each compound. Predicted pChEMBL values are annotated only for the HEV and LEV compounds in the first row because they were allocated to the test set. The nearest neighbors (rows 2–4) were allocated to the training set and, therefore, no prediction of the pChEMBL value was made during evaluation. Left panel: an activity cliff example with the HEV compound in the first row and its three nearest neighbors below. Right panel: LEV compound in the first row and its three nearest neighbors below.

CV and the results are shown in SI Table 5. Targets that yielded negative  $Q^2$  scores across all three models were excluded from the comparative analysis. The remaining 31 targets that were considered for the aggregate analysis comprised 8698 unique compounds, representing 95.4% of the dataset. When compared to other methods in Table 8, the MT-DNN model outperformed the single-task models in all categories, except for the aggregate standard deviation (SD) scores of the ST-RF model in the unweighted and weighted categories. The most notable improvement was observed in the inversely weighted category, indicating that underrepresented targets mostly benefited from the MT-DNN model without compromising performance in other categories. Additionally, more than 60% of the targets showed improved scores with the multi-task learning (MTL) approach.

To showcase discrepancies between positive and negative transfers with MTL, three specific targets with different

performance profiles were analyzed using a 2D representation of the chemical space in Fig. 5. The entire dataset was included in the chemical space analysis to represent relationships between individual clusters. Starting with the highest-scoring target in Table 9, SLC5A1 (sodium/glucose co-transporter, SGLT1) had a  $Q^2$  of 0.809. The chemical space of compounds associated with SLC5A1 shows a wider degree of coverage between multiple domains when compared to the other two targets in Fig. 5. The compounds of the lowest-scoring target, SLC33A1 (acetyl-coenzyme A transporter 1), with a  $Q^2$  of  $-2.554$ , contrast with the distribution pattern of the SLC5A1 chemical space by having a localized and entirely isolated cluster. Based on this representation, it is evident that no other member shares similar chemical space with SLC33A1. It cannot be excluded that the cause of the discrepancy in the performance between the two targets stems from quantitative imbalance of the dataset. SLC5A1 contributes 975 datapoints to the overall



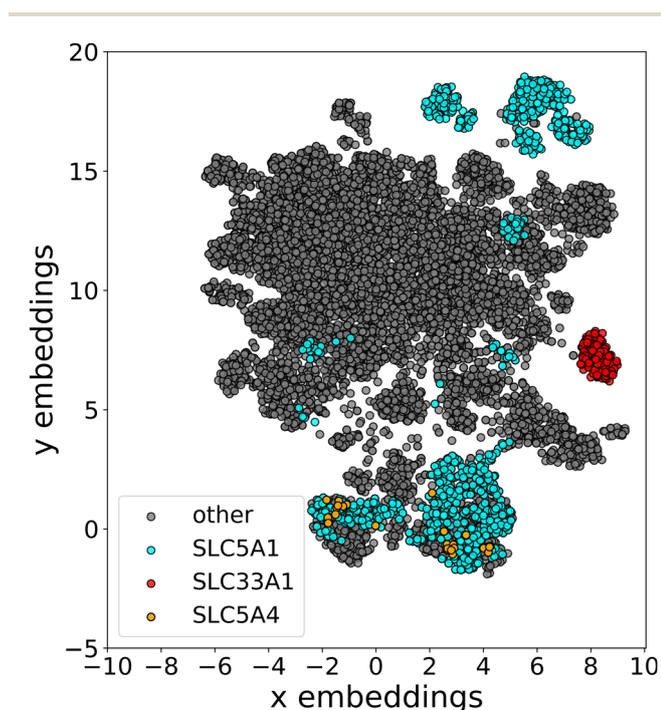
**Table 8** Target-based aggregate results for MT-DNN, ST-DNN, and ST-RF models under three weighting schemes: unweighted (equal target contribution), weighted (proportional to the number of compounds), and inversely weighted (emphasizing smaller targets with an inversely proportional number of compounds)<sup>a</sup>

Model	Unweighted			Weighted			Inversely weighted		
	$Q^2$	$Q^2 - SD$	TSH (%)	$Q^2$	$Q^2 - SD$	TSH (%)	$Q^2$	$Q^2 - SD$	TSH (%)
MT-DNN	<b>0.280</b>	0.267	<b>61.290</b>	<b>0.521</b>	0.098	<b>51.556</b>	<b>0.247</b>	<b>0.326</b>	<b>70.770</b>
ST-DNN	-0.050	0.590	12.903	0.454	0.169	33.026	-0.475	1.046	5.299
ST-RF	0.232	<b>0.205</b>	25.806	0.496	<b>0.075</b>	15.417	0.069	0.340	23.931

<sup>a</sup> Every category has three scores across 31 targets:  $Q^2$  – mean of cross-validated coefficients of determination,  $Q^2 - SD$  – mean of standard deviations for the  $Q^2$  value, and TSH – percentage of targets that improved with the corresponding model.

dataset, whereas SLC33A1 contributes 131 datapoints. Furthermore, SLC5A1 belongs to the SLC5 family with three additional protein members modeled as separate tasks that collectively amount to 2194 compounds in the dataset. Such instances of correlated tasks would, as stated in the introduction, be able to leverage the shared representation of knowledge

generated with an MTL approach. To further support this hypothesis, a  $Q^2$  value of 0.758 is observed for the SLC5A4 (sodium/glucose cotransporter 3, SGLT3) target with 28 compounds in the dataset. Although it was represented by fewer datapoints, SLC5A4 scored higher than SLC33A1 in Table 9. As highlighted in Fig. 5, compounds of SLC5A4 share the chemical space with other targets and do not form isolated clusters. Unlike SLC33A1, which was the single member of the SLC33 family in the dataset, SLC5A4 belongs to the SLC5 family that is shared with three other members. Furthermore, comparison of the performance scores between the MT-DNN and ST-RF models in Table 9 suggests that the SLC5 members benefit from the MTL approach, whereas the SLC33A1 target was able to achieve a higher score with the ST-RF model. Understudied targets such as SLC5A4 have little information published on their active compounds when compared to some other members of the SLC superfamily. To the best of our knowledge, this MTL model presents itself currently as a unique tool for compound bioactivity prediction and imputation of the SGLT3 (SLC5A4) transporter.



**Fig. 5** Chemical space representation with UMAP. Individual compounds are colored according to associated targets marked in the legend.

#### Data imputation performance

An imputation-dedicated split was chosen to evaluate the ability of an MTL approach to predict the pChEMBL values of compounds for targets not used in the output layer during training. This means that the model is tested on compounds that are available in the training set but show different output profiles. Unlike the conventional approach where the model is tested on an entirely new and previously unseen set of compounds, this step examines how well the model performs in an imputation-driven setting by estimating pChEMBL values of known compounds for a different target.

**Table 9** Target-based evaluation with three examples highlighted for each performance case<sup>a</sup>

Target	Compounds	Family members	Total compounds	MT-DNN ( $Q^2$ ) <sup>↑</sup>	ST-DNN ( $Q^2$ ) <sup>↑</sup>	ST-RF ( $Q^2$ ) <sup>↑</sup>
SLC5A1	975	4	2194	<b>0.809 ± 0.006</b>	0.795 ± 0.021	0.799 ± 0.025
SLC33A1	131	1	131	-2.554 ± 1.548	-2.866 ± 3.151	<b>0.086 ± 0.198</b>
SLC5A4	28	4	2194	<b>0.758 ± 0.107</b>	0.428 ± 0.587	0.667 ± 0.201

<sup>a</sup> The table shows the quantitative properties of selected targets with individual performance scores represented as  $Q^2$  and standard deviation from the MT-DNN, ST-DNN, and ST-RF models. The table additionally shows the number of compounds for each target and their respective families. Bold – the best value in a row between the three models.



**Table 10** Comparison between optimized parameters of the prediction and imputation split models

Hyperparameters	Models	
	CDDD prediction	CDDD imputation
Input size	512	512
H1 size	800	1020
H2 size	250	170
H3 size	—	80
Task-specific size	54	54
Learning rate	0.000043	0.000105
Dropout rate	0.1	0.09
Patience	10	10

Since neither the CDDD nor ECFP6 prediction models achieved a minimum MSE validation value with the inclusion of task-specific neural network domains (Table 5), only the shared domain was incorporated as a parameter during the optimization of the CDDD imputation model. As shown in Table 10, the CDDD imputation model achieved the lowest MSE value for the validation set with higher architectural complexity than the prediction counterpart. One additional hidden layer was selected along with a higher learning rate and identical patience to counter overfitting.

The CV step was excluded in this section due to the lack of a sufficient number of compounds with five or more bioactivity measurements for accurate allocation in each fold. The results in Table 11 show that CDDD imputation was able to score relatively close to the CDDD prediction model. A slight improvement in the prediction performance over the imputation performance could be attributed to the skewed data distribution introduced by the imputation split, which may affect generalization. The imputation split only considers compounds with two or more measurements, potentially biasing target representation, molecule types, and activity values.

Comparative analysis between accuracy scores on the test set in Table 12 shows that the MT-DNN outperforms all six imputation methods chosen for benchmarking. The second-highest

**Table 12** Benchmarking MT-DNN performance against six different imputation methods<sup>a</sup>

	sklearn.imputers				fancyimpute		
	MT-DNN	S-mean	KNN	Iterative	Soft	I-SVD	MF
$R^2 \uparrow$	<b>0.720</b>	0.098	0.672	0.471	-0.400	0.107	0.489
MSE $\downarrow$	<b>0.407</b>	1.312	0.477	0.667	2.033	1.299	0.744

scoring method is k-Nearest Neighbor from the sklearn.imputers module. Considering both high prediction and imputation performance, MTL presents itself as a suitable and versatile tool in bioactivity-value estimation of compounds towards SLC transporters.

## Conclusion

With this study, we present an MTL-based prediction to complete a bioactivity matrix of compounds for the superfamily of SLC transporters. Besides replacing the missing values, the MT-DNN could predict bioactivity profiles for out-of-sample compounds and showcase overall satisfactory performance scores. Further results indicate that the selection of different types of descriptors can affect the ability of the model to achieve reliable predictions. In this case, the causality of descriptor-type discrepancies in the performance is not completely understood, and further studies are necessary for conclusive statements. Based on individual bioactivity-prediction error analysis and target-based evaluation, it cannot be excluded that the shortcomings of our MTL approach are associated with the presence of activity cliffs and heterogeneity between the modeled targets. The dataset contains compounds with similar structural properties but contrasting bioactivity values or sets of associated targets. If imbalanced in quantity and distribution between the training, validation, and test sets, such outlier instances designated as activity cliffs can impact the performance of the model. A similar phenomenon was observed for a target-specific outlier, where the MTL model struggled to accurately predict the bioactivity values of compounds that formed an isolated cluster in the chemical space analysis. Conversely, the performance of SLC5A4 suggests that a target with a low quantity of datapoints could benefit from the MTL approach if it shares certain qualitative and quantitative properties with other correlated targets, such as having common families or similar chemical space. In conclusion, the comparative analysis demonstrates the suitability of the multi-task model as a prediction tool intended for a data imputation task. Besides scoring highest during benchmarking against six established imputation methods, MT-DNN outperformed the single-task models by achieving higher accuracy for underrepresented targets. Despite this improvement bias, the MT-DNN model

**Table 11** Comparison between the scoring metrics of prediction and imputation assessment<sup>a</sup>

	Metrics	Models	
		CDDD prediction	CDDD imputation
Validation set	MTS $\uparrow$	<b>0.942</b>	0.940
	$R^2 \uparrow$	<b>0.754</b>	0.705
	MSE $\downarrow$	<b>0.377</b>	0.429
	MAE $\downarrow$	<b>0.465</b>	0.503
Test set	RMSE $\downarrow$	<b>0.615</b>	0.655
	$R^2 \uparrow$	<b>0.739</b>	0.720
	MSE $\downarrow$	<b>0.362</b>	0.407
	MAE $\downarrow$	<b>0.451</b>	0.477
	RMSE $\downarrow$	<b>0.602</b>	0.638

<sup>a</sup> Bold – the best value in a row.



achieved comparable or slightly higher scores for most other targets with larger numbers of compounds, highlighting its strengths in handling missing bioactivity data for an imbalanced target space.

## Author contributions

TC: conceptualization, data acquisition, data processing and standardization, literature research, methodology, evaluation, visualization and validation of the results, Python code writing and documentation, and writing of the manuscript. SS: conceptualization, supervision, and draft review. GFE: conceptualization, project administration, supervision, resources, and draft review.

## Conflicts of interest

There are no conflicts of interest to declare.

## Data availability

The data used for this study are available on the Resolute Knowledgebase at: <https://doi.org/10.5281/zenodo.4309586>, <https://re-solute.eu/knowledgebase/gene>. The UniProt dataset is available at: <https://doi.org/10.1093/nar/gkae1010>, [https://www.uniprot.org/uniprotkb?query=\\*%&facets=model\\_organism%3A9606](https://www.uniprot.org/uniprotkb?query=*%&facets=model_organism%3A9606). The intermediary datasets generated after standardization, processing, and calculating descriptors are available on the GitHub repository along with the Jupyter Notebook and Python scripts used for the development of this study at: <https://doi.org/10.5281/zenodo.17991114>, <https://github.com/PharminfoVienna/SLC-data-imputation>.

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